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- (54) EX-VIVO ISOLATED CD25+CD4+ T CELLS WITH IMMUNOSUPPRESSIVE ACTIVITY AND USES THEREOF

EX-VIVO ISOLIERTE CD25+CD4+TZELLEN MIT IMMUNSUPPRESSIVER AKTIVITÄT UND DEREN ANWENDLINGEN

LYMPHOCYTES T CD25+CD4+ ISOLES EX VIVO A ACTIVITE IMMUNOSUPPRESSIVE ET UTILISATIONS

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Description

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[0001] The present invention provides ex-vivo isolated human C025*C04* T regulatory (Tr) cells, homogeneous clonal populations derived therefrom with enhanced suppressive activity and their uses in the regulation of immune responses and for the identification and characterization of suppressor T cell specific molecules. More specifically, the invention is directed to the use of polyclonal CD25*C024* Trotals or of homogeneous clonal C025*C024* Trotals generated ex-vivo to prevent or treat conditions where a down-regulation/suppression of the immune response is required, such as graft-vs-host disease (GVHD), organ rejection, gene therapy and autoimmune diseases, or for the identification and characterization of molecules involved in the regulation of the immune response is reconstitution.

BACKGROUND OF THE INVENTION

[0002] T regulatory (Tr) cells have a key role in the maintenance of immune tolerance to both self and harmless foreign antigens. Many subsets of Tr cells have been described and recently much progress has been made in understanding their ontogeny, function and mechanisms of action (reviewed in (1)). Some Trcells do not produce cybidness and suppress T-cell responses via a mechanism that requires direct cell-cell contact (2, 3). Other subsets of Tr cells produce immunoregulatory cytokines, such as IL-10 and TGF-β, and exert their suppressive functions at least in part via the effects of these crybidness (4-8).

[0033] CD4* Tr calls that constitutively express the IL-2R chain (CD25) have been identified in the mouse (reviewed in (2, 3)). These CD25*CD4* Tr calls show a remarkable suppressive capacity both in vitro and in vivo. Transfer of these Tr calls reduces the pathology of experimentally-induced autoimmune diseases such as thyroidfist, gastrills, insufrindependent cliabetes mellitus and collist (6+12) and of experimentally induced GvHD (31), whereas depiction of CD25*CD4** Tr cells results in the development of vistemic autoimmune diseases (11, 13, 14).

[0004] Murine CD25*CD4* Tr cells are anergic when stimulated in vitro with anti-CD3 mAbs, but proliferate upon addition of soxponous IL-2 (16, 16). After TCR-mediated stimulation, CD25*CD4* Tr cells suppress the activation and proliferation of other CD4* and CD8* T cells in an antigen non-specific manner (16, 17) via a mechanism that requires cell-cell contact and that, in most systems, is independent of production of immunosuppressive cyclokines (15, 16). Murine CD25*CD4* Trcells constitutively express cytokovic Tymphocyte-associated antigen at (CTLA-4)(9), a negative regulator of T-cell activation, and expression of this molecule is required for the ability of these cells to suppress immune responses in vivo (10, 18). In addition, CD25*CD4* Tr cells may act by down-regulating the expression of CD0 and CD86 on APCs (19), although some reports suggest that APCs are not required for their suppressive activity and indicate that direct T-T cell interaction is involved (17).

DESCRIPTION OF THE INVENTION

[0005] This invention is based on the findings that human CD25*CD4* Tr cells with immunosuppressive effects can be isolated from peripheral blood and expaned for vitro without loss of function, and that human CD25*CD4* Tr cells constitute a heterogenous population from which different cell clones exhibiting suppressive or non-suppressive activity can be derived and solated based on expression of CD25. Isolated human CD25*CD4* Tr cells and CD25*CD4* To cell clones can be used as immunosuppressive agents for the prevention or treatment of pathologies where a reduction of the immune response is desired. Typically, they will be used to prevent GH2D, organ rejection, immune responses directed against foreign proteins introduced during gene therapy and autoimmune diseases, especially type 1 diabetes. CD25*CD4* Tr cells isolated from peripheral blood can be stimulated and cultured in vitro, allowing for the possibility to select and expand entiplen-specific suppressor Toals. Expanded CD25*CD4* Troels is maintain their regulatory capacity and vitro, and thus can be used to regulate T cell responses in vitro, whereas both freshly-isolated and in vitro-expanded human CD25*CD4* Troells can be utilized in therappin vivor. The methods and conditions for isolation and in vitro expansion of CD25*CD4* Troells can be expanded in vitro under one or more of the following conditions: co-culture with feeder-cell mixture, polydonal stimulation, antigen specific stimulation, addition of cytokines.

[0006] The T cells thus obtained can be re-introduced in the patient. The preferred modalities under which CD25*CD4*

Tricells are used in therapy or prophylaxis will depend on the particular condition to prevent/treat.

[0007] For example, to prevent/real GvHD, CD25*CD4* Tr can be isolated from leukapheresis of the bone-marrow donor, frozen if necessary, and administered to the recipient at the time of transplant, prior to the transplant or in the subsequent months. Alternatively, CD25*CD4* Tr cells from the donor could be stimulated with host APC in vitro, in order to generate and expand alloantigen-specific CD25*CD4* Tr cells that would specifically suppress host-specific responses in vitro.

[0008] To prevent/treat organ rejection, CD25+CD4+ Tr cells can be isolated from the recipient, frozen if necessary, and administered prior to transplant, at the time of transplant or in the subsequent months. Alternatively, CD25+CD4+

Tr cells could be stimulated in vitro with autologous APCs that have been or-cultured with tissue from the organ in question and will therefore present foreign antigens trough the indirect pathway (32). The resulting CD25°CD4 Tr cell lines would be specific for antigens expressed by the transplanted organ and could be used to suppress organ-specific

[0009] To prevent autoimmune diseases, bulk populations of autologous CD25°CD4* Tr cells can be isolated and reinfused. Alternatively, antigen-specific CD25°CD4* Tr cells could be expanded in vitro by stimulation with autologous APCs and self-antigens derived from tissues which are targets of the disease. Upon re-administration of the in vitroexpanded CD25°CD4* Tr cells, they will suppress anti-self responses in vivro.

[0010] To prevent immune response in gene therapy, CD25°CD4° Tr cells can be isolated from the recipient, and to cells which are specific for entigens encoded by the therapeutic vector could be expanded in vitro by stimulation with transduced autologous APCs expressing the transgene. These cells can be frozen if necessary, and administered at the time of cene therapy treatment or in the subsequent months.

[0011] Advantageously, a homogeneous clonal population of CD25°CD4° Tr suppressive cells is used for the therapeutic applications indicated above. The method for isolating suppressive CD25°CD4° Tr clones essentially comprises the stens of:

a) purifying CD4+ T cells from PBMCs;

b) separating CD25+ from CD25- T cells;

c) cloning CD25+CD4+ T cells by limiting dilution:

d) stimulating with phytohemagglutinin (PHA) or anti-CD3 mAb in the presence of IL-2;

e) selecting the cell clones that display a constitutively high expression of CD25.

[0012] According to step (a), CD4+ cells can be purified by positive selection with anti-CD4-coupled microbeads. Step (b) can be carried out by marking CD25+ cells using labelled anti-CD4/25 monoclonal antibodies and purifying CD25+ cells by FAC5-sorting. The clones obtained from step (c), which can be maintained in X-vivo 15 culture-medium or in other cellular media, supplemented with 5% pooled or autologous human serum, are preferably restimulated by couture with feeder-cell mixture, by antigens or by cytokines, more preferably by an allogenio or autologous feeder-cell mixture consisting of irradiated PBMCs, with or without irradiated autologous or allogeneic EBV-transformed cell lines (eg. UT). Suppressive clones which display a constitutively high expression of CD25 can be selected, according to step (d), on the basis of the following characteristics:

- 100% constant-positivity for CD25 expression in the resting phase at least 10 days after stimulation with phytohemagglutinin or anti-CD3 mAb in the presence of IL-2;
- expression of CD25 at a significantly higher level in comparison to T cell clones isolated in parallel from CD25 CD4+
 T cells or non suppressive clones isolated from CD25+CD4+T cells.

[0013] At the end of steps (a)-(d), homogeneous CD25*CD4* T-cell clones constitutively expressing CD25, anergic and with high suppressive capacity are isolated. The suppressive clones, in contrast to the non-suppressive ones, are characterized by significant production of TGF-B and no production of IL-2.

40 [0014] In a further embodiment, the invention provides an immunosuppressive agent containing isolated CD25*CD4*Tr-cells and/or CD25*CD4*Tr cell clones constitutively expressing CD25, and optionally other active substances, such as cytokines, or other immunosuppressive proteins. Preferably the immunosuppressive agent will be in the form of a stabilized cell preparation.

[0015] Besides the envisaged clinical applications, the CD25*CD4* Tr suppressive cell clones can be used to set up systems in vitro for the identification of molecules that modulate the immune response, in particular suppressor. T-cellspecific molecules. According to a preferred embodiment, such CD25*CD4* Tr cell clones will be used in large scale gene expression arrays, in differential proteomics screenings and in the generation of monoclonal antibodies specific for Tr cells constitutively expressing CD25. These applications are greatly aided by the homogeneity of comparative samples, such as that provided by cell populations of clonal origin.

DESCRIPTION OF THE FIGURES

[0016]

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Figure 1. Isolation and cell-surface phenotype of human CD25°CD4* Tr. cells: CD4* To cells were solated from FBMCs, and separated into CD25* and CD25* retactions. Purily (A) and expression of CD45*CD, HLAP (B), IL-2R), and CD824 (C) was determined by FACS. CD25*CD4* and CD25*CD4* To cells were either cultured in medium ations or activated with immobilized anti-CD3 mAbs or PMA and calcium immophore for 6 hours (D) or 24

hours (E). Cells were analyzed for surface-expression of CD40L and CD69 (D), and for intracytoplasmic expression of CTLA-4 (E). Results are representative of 6 independent experiments.

Figure 2. CD25*CD4*T reals are amergic and suppress proliferation to alloantiques. Purified CD25*CD4*T reals (10,000 cellavell) were tested for their ability to proliferate in response to immobilized ant-CD3 mAbs (aCD3) (10µg/ml) in the absence or presence of soluble anti-CD28 mAbs (aCD28) (1µg/ml), secondary rabbit anti-mouse Abs (aCD28*CL) (10µg/ml), and/ort.-2 (100U/ml). After 72 hours of culture, 91-thymidine was added for an additional in hours (a). CD25*CD4*T cells (6), COD25*CD4*D4*C cella diagonaic APCs in the absence or presence of increasing numbers of autologous CD25*CD4*T cells (7), CD25*CD4*T cells were activated to induce CD25*expression. After 48 hours T cells that became CD25**CD25*D4*T cells were activated to induce CD25*expression. After 48 hours T cells that became CD25**CD25*D4*T cells were activated by alloantigens with or without CD25*CD4*T r cells in response to alloantigens (C), CD25*CD4*T cells were activated by alloantigens with or without CD25*CD4*T r cells in response to alloantigens (C), CD25*CD4*T cells were activated by alloantigens with or without CD25*CD4*T r cells (7), in the presence of the indicated mAbs continued to the continued of the continued to the cont

Figure 3. Expansion and cell-surface phenotype of CD25*CD4* Tr cells. CD25* and CD25*CD4* T cells were purified, and activated with anti-CD3 mAbs, allogeneb feeder-cell mixture and exogenous IL-2. Cells were split as necessary and after 2 weeks were analyzed by CR56 for expression of CD25 and CD4 (A). In parallel, cells were analyzed for cell-surface expression of CD40 and CD69 following activation for 6 hours with immobilized anti-CD3 mAbs or PMA and calcium lonophore (B). Constitutive levels of CTLA-4 expression was determined by intracyto-plasmic staining (C). Results are representative of 3 independent experiments.

Figure 4. Cultured CD25*CD4* Tr cells retain their suppressive capacity. CD25* and CD25*CD4* Tr cells were purified and activated with anti-CD3 mAbs, allogeneic feeder cell mixture and exogenous IL-2. After 14 days of culture, T cells were tested for their ability to proliferate in response to anti-CD3 mAbs (10μg/ml) in the absence or presence of soluble anti-CD26 mAbs (10μg/ml) and/or IL-2 (100U/ml) (A). Cultured CD25 CD4* To cells (6,0). CD4* coll (6,0). CD4

Figure 5. Isolation of human CD25*CD4* T cells at the clonal level. CD4* T cells were isolated from peripheral blood, stained with anti-CD4 and cm1-CD25 mAbs, and separated into CD25*CD4* and CD25*CD4* T cells by FACS sorting to a purity greater than 98 and 99% respectively.

Figure 6. CD25*CD4* T cell clones are heterogenous in terms of their expression of CD25 in the resting phase. Resting T-cell clones were stained with anti-CD24 and -CD25 mAbs 12-14 days after the last re-stimulation, The number (#) of the T cell clone is indicated on the upper left, and the MFI and percent of CD25 positive cells is on the upper right.

Figure 7. CD25*CD4* T cell clones are heterogenous in term of their proliferative response to activation via the TOR Insting T-cell clones were tested for their ability to proliferate in response to anti-CD3 mAbs (10µg/m) in the 50 cases of the proper of the 2 (100µm). After 48 hours of culture, 94-thymridine was added for an additional 16 hours. Figure 8. Suppression of naive 7 cell responses by CD25*CD4*T cell clones. Autologous CD4*T cells were purified and tested for their ability to proliferate in response to anti-CD3 mAbs and irradiated CD3-depleted APCs (A) or anti-CD3 mAbs immobilized on plastic (B). After 72 (A) or 48 hours (B) of culture, 94-thymridine was added for an additional 16 hours. Numbers indicate percent reduction in proliferation in comparison to the naive CD4*T cells atone.

DETAILED DESCRIPTION OF THE INVENTION

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Isolation and cell-surface phenotype of human CD25+CD4+ Tr cells

[0017] CD25*CD4* Trcells are present in human PBMCs. On average they represent 3.0% (range 1.6-4.4%, n.=5) of total PBMCs and 12.8% (range 9.8-18.1 %, n.=5) of CD4* T cells. These cells could be readily losteded, with purities greater than 90% (Figure 1A). The majority (82 ± 5.1%) of freshly isolated CD25*CD4* Trcells also expressed CD45*D0 and the percentage of CD25*CD4* Tr cells expressing HLA-DR was significantly higher than that in the CD25 CD4* population (17.3 ± 4.9% vs 64 ± 2.9%, n.=6) (Figure 1B). In addition, human CD25*CD4* Tr cells expressed dipher levels of the IL-2Rβ in comparison to CD25*CD4* T cells (19.9 ± 2.6% vs 93.1 ± 2.5%, n.=5) (Figure 1C) and a subset of freshly isolated CD25*CD4* Tr cells expressed CTLA-4 (8.4 ± 1.6%, n.=5) (Figure 1E). In contrast, expression of the IL-2Rh and CD25*CD4* Tr cells expressed of CD25*CD4* Tr cells and CD25*CD4* To cells were

CD3+TCR α β +, Thus, human CD25+CD4+ Tr cells express markers which are characteristic of memory T cells and low constitutive levels of CTLA-4, similarly to murine CD25+CD4+ Tr cells (9, 10, 15, 17, 18).

[0018] Following TCR-mediated stimulation, human CD25*CD4*T recells expressed lower levels of activation markers in comparison to CD25*CD4*T recells. The proportion of CD40L-0 tsless are 1.73 = 2.9% in CD25*CD4*T recells vs 28.4 = 1.8% in the CD25*CD4*T-cells ubset (p 0.005); whereas 30.9 ± 7.0% of CD25*CD4*T recells vs 54 ± 9.1% of CD25*CD4*T cells expressed CD59 (p 0.006); [Vignor ID). Time course experiments demonstrated that the reduced levels of CD40L and CD69 on CD25*CD4*T recells was the very constraint of the reduced levels of CD40L and CD69 on CD25*CD4*T recells were not due to eithered kinetics of expression. After activation with PMA and calculum ionophore there was no statistically significant difference between expression of CD69 or CD40L on CD25*CD4*T or CD25*CD4*T cells, although in general fewer CD25*CD4*T recells, although in general fewer CD25*CD4*T recells, although in general fewer CD25*CD4*T recells although in general fewer CD25*C

[0019] Following activation with anti-CD3 mAbs or PMA and calcium inorphore, the percentage of CD25*CD4* Tr. cells expressing CTLA-4 was higher than that of CD25*CD4* T cells. In addition, CTLA-4 expression levels were approximately 3 fold higher on CD25*CD4* Tr. cells (Figure 1E). Collectively, these data demonstrate that upon TCR activation human CD25*CD4* Tr. cells have a surface molecule expression profile which is unique and distinct from that of other CD4*T-cell subsets.

Proliferation and cytokine production by human CD25+CD4+ Tr cells

alloantigen stimulation than did CD25CD4+ T cells.

[0020] Freshly isolated CD25*CD4*Trcels did not profilerate in response to immobilized anti-CD2 mAbs, and addition of soluble anti-CD28 mAbs resulted in a modest and variable increase in profileration. In contrast, crosslinked anti-CD28 mAbs completely reversed the unresponsiveness of CD25*CD4*Tr cells to TCR addition of exogenous L-2 partially restored the profileration of CD25*CD4*Tr cells in response to anti-CD3 mAbs, and profileration was further enhanced by soluble anti-CD28 mAbs Figure 2A). These results indicate that human CD25*CD4*Tr cells have a specific defect in their ability to profilerate after TCR-mediated activation (15.

[0021] Human CD25*CD4* Tr celle were analyzed for their ability to produce cytokines. Following stimulation with immobilized antri-CD3 mAbs, with or without soluble anti-CD28 mAbs, no detectable levels of IL-2, IL-10, IL-4, TGF-jb or IFN-y could be measured. In contrast, when stimulated with anti-CD3 and soluble anti-CD28 mAbs in the presence of exogenous IL-2, CD25*CD4* Tr cells produced significant levels of IL-4, IL-10, IFN-y and TGF-jb (Table 1). Under these stimulation contraines CD25*CD4* Tr cells ind a cytokine profile that was comparable to that of CD25*CD4* T cells in ontrast, differences in cytokine production were observed following activation with allogenes APCs. Both CD25*CD4* and CD25*CD4* T cells produced IL-10, TGF-jb and IFN-y, but no IL-1. However, the striking difference between the CD25*CD4* and CD25*CD4* T cell populations is that the CD25* cells failed to secrete IL-2, indicating that these cells wave a specific detect in production of IL-2. Interestingly, CD25*CD4* T r cells considerative produced less FIN-y upon

5 Human CD25+CD4+ Tr cells suppress the proliferative responses of naive CD25-CD4+ T cells to alleantigens.

[0022] We investigated the regulatory properties of CD25*CD4** Troells by testing their ability to suppress the proliferative responses of naive CD25*CD4** To ells to ability 10s suppress the proliferation of acceptance of the control of the cont

[0023] In order to demonstrate that this suppressive capacity was an intrinsic property of T cells constitutively expressing DDEs in vivo, we tested whether CDDS CD4 To ells which expressed CD25 following activation in vitrosowed regulatory effects. To this aim, CD25 CD4* T cells were activated with arti-CD3 and arti-CD2 amAbs, and after 48 hours the CD25* T cells were isolated and tested for their ability to suppress freshly isolated autologous CD25* Tcells. As shown in Figure 2C, T cells which became CD25* that in vitro activation proliferated in response to alloantignes, and enhanced rather than suppressed the response of CD25* CD4* T cells. These data indicate that inhibition of T cell proliferation is a unique property of cells which constitutively express CD25* in vivo.

50 [0024] Several studies show that some subsets of IT cells, such as type 1 Tr (Tr1) and Th3 cells, produce IL-10 and/or TGF-β and suppress immune responses via production of these cytokines (4.8). Since CD25*CD4* Tr cells produced both IL-10 and TGF-β upon stimulation with allogeneix APCs (Table 1) the role of these cytokines on inhibition of allogeneir responses by human CD25*CD4* Tr cells was investigated. As shown in Figure 2D, addition of neutralizine.

anti-IL-10R or anti-TGF- β mAbs had no significant effect on the ability of CD25*CD4* Tr cells to suppress the proliferation of CD25*CD4* To cells in response to alloantiques. In 3 independent experiments an average of 572 \pm 5.0% suppression was observed in the presence of 10 μ g/mil of control 19 $_{\rm G}$ 6.83 \pm 5.8% with 10 $_{\rm G}$ 9ml of mil-IL-10R and 80.2 \pm 3.4% with 10 $_{\rm G}$ 9ml of anti-TGF- β mAbs. Similar results were obtained with a 5 fold higher concentration of anti-TGF- β mAbs. Addition of both arti-IL-10R and anti-TGF- β mAbs resulted in a slight, but not statistically significant, reversal of suppression mediated by CD25*CD4*T crolls from 872 \pm 2.6% of 562 \pm 5.7%, b, 0.06.

[0026] F(ab), fragments from antibodies which specifically block the ability of CTLA-4 to bind to CD80/88, without affecting eignals via CD28, have previously been shown to inhibit the production of TGF-β by Trl cells (20). Addition of the same blocking anti-CTLA-4 mAbs had no significant effect on the suppressive activity of CD28* CD4* Tr cells. These data suggest that despite the fact that CD28* CD4* Tr cells express high levels of CTLA-4 (Figure 1E), this molecule is not required for their suppressive activity.

Human CD25+CD4+ Tr cells can be expanded in vitro

19 [0026] We have previously shown that human Tr1 cells which have been cloned and expanded in vitro maintain their regulatory activity (6). Using a protocol similar to that described for Tr1 cells, we determined whether CD25+CD4+ Tr cells cid not proliferate in response to anti-CD3 slone (Figure 2A), but when activated with anti-CD3 mAbs in the presence of an allogene's feeder-cell mixture and exogenous IL-2, an expansion which was entillar to that of CD25-CD4+ T cells was obtained (20-30 fold increase at day 14). In vitro-expanded human CD25+CD4+ T cells remained positive for CD25 even after culture for more than one month (Figure 3A). In contrast, all CD25-CD4+ T cells expressed CD25 after activation, but the expression gradually decreased with time. Persistent expression of CD25 has also been observed in myrine CD25-CD4+ T cells activated in vivo (21).

[0027] Similar to freshly isolated CD25*CD4* Tr cells, the proportion of *in vitro*-expanded CD25*CD4* Tr cells expressing CD40. I following activation with anti-CD3 mAbs was consistently lower in comparison to CD25 CD4* To cells (2.7 ± 3.3% vs 9.5 ± 8.3%, p.0 to). However, incontrast to freshly isolated cells, cultured CD25*CD4*Tr cells expressed normal levels of CD89 following polyclonal TC4-mediated activation (Figure 38). Thus, reduced up-equilation of CD40, is also a characteristic of expanded CD25*CD4* Tr cells. As expected, cultured CD25*CD4* and CD25*CD4* To cells expressed similar levels of both CD40, and CD89 following activation with PMA and calcium inorphore. *In vitro*-expansion of CD25*CD4* Tr cells of the contribution of CD40, and CD89 following activation with PMA and calcium inorphore. *In vitro*-expansion of CD25*CD4*T cells expressed similar developed to express this inhibitory molecule at sindifficantly higher levels than the CD25*CD4* contributants (Flour 9.3).

In vitro-expanded CD25*CD4* Tr cells remain anergic and retain their suppressive capacity

45 [0029] In the present study we show that human CD4+ T cells which express CD26 in who are a unique subset of Tr cells. Human CD25+CD4+ T reclis are anergic, fall to produce IL-2, constitutively express CTLA-4, and suppress the proliferation of neive CD4+ T cells, as described for murine CD25+CD4+ Tr cells (2, 3). In addition, following polycional TCR-mediated activation, human CD25+CD4+ Tr cells strongly upregulate CTLA-4, display reduced express on of CD40L, and produce cytokines. CD25 and CTLA-4 remain constitutively expressed on in intro-expanded human CD25+CD4+ To cells, while following activation, up-regulation of CD40L is still defective. More interestingly, in vitro expanded CD25+CD4+ T cells reach their potent suppressive activity, even towards previously additated memory T cells. The observation that functional Tr cells can be expanded in vitro and can regulate responses of memory T cells is of great clinical relevance for the use of CD25+CD4+ T cells as a cellular therapy in T-cell mediated diseases.

[0030] The role of Immunoregulatory-cytokines in the suppression mediated by CD25*CD4* Tr cells remains an open question. Alloantigen-activated CD25*CD4* Tr cells did not proliferate but produced IL-10, IFN-y-and TGF-β, and indeed possessed a profile of cytokine production which is comparable to that of Tr1 cells (i.e. IL-10+IFN-y-TGF-β-IL-L-IL-2-I/ov) (6). However, we observed only a slight reversal of suppression in the presence of both neutralizing anti-IL-10A and anti-TGF-3 mAbs. which is consistent with previous observations that production of IL-10 and TGF-8 is dissensable.

for the regulatory function of CD25°CD4° Tr cells (15, 16). On the other hand, in a murine model of experimentallyinduced colitis, both IL-10 and TGF-β were found to be required for suppression mediated by CD25°CD4° Tr cells (5, 10). The basis for this discrepancy in the involvement of IL-10 and TGF-β is unclear. CD25°CD4° Tr cells have the capacity to produce IL-10 and TGF-β, but production of these cytokines may depend on their maturation state and the environmental contract in which they are activated.

[0031] Previous reports demonstrated that direct cell-cell contact is required for murine CD25*CD4*Tr cells to exert their suppressive effocts (15, 16). Despite constitutive and persistent expression of CTLA4, anti-CTLA4 mbAs failed to abrogate the suppressive activity of human CD25*CD4*Tr cells. These data are in agreement with a study indicating that signals via CTLA4 were dispensable for suppression by mouse CD25*CD4*Tr cells in vitro (15). However, more recent reports indicate that expression of CTLA4 is essential for suppression mediated by these cells (10, 18), it is possible that suppression of profiteration operates via mechanisms which differ depending on the stimuli and microenvironment, or that human and mouse CD25*CD4*Tr cells actitivough different mechanisms. Finally, suppression is not simply due to consumption of It.2 as murine CD25*CD4*Tr cells suppressed It.2-production at the transcriptional level (15). In addition, human CD4*T cells which expressed CD25*CD4*Dr on simply result in sequestration of It.2.2.

[0032] Human CD25*CD4* Treelis expand in vitro and maintain their unique cell-surface marker profile and suppressive functions. To our knowledge, these data represent the first report of in vitro expansion of human T suppressor cell lines

[0033] The clinical use of CD25rCD4+ T regulatory (Tr) cells can be envisaged to down-regulate undesired immune responses in a number of pathological conditions. We have shown that CD25rCD4+ Tr cells with suppressive function can be readily isolated from peripheral blood. Importantly, these cells can be stimulated and cultured in vitro, allowing for the possibility to select and expand antigen-specific suppressor T cells. Expanded CD25rCD4+ Tr cells meintain their regulatory capacity in vitro, and thus could be used to regulate T cell responses in the CB1 reports in Vitro.

Isolation and characterization at the clonal level of human CD25+CD4+ T cells with suppressive capacity

[0034] In order to determine the relationship between IL-10 producing Tr1 cells (22) and CD25*CD4* Tr cells (23, 24), we attempted to isolate CD25*CD4* Tr cells at the clonal level. It has previously been described that only approximately 0.8% of peripheral blood monouncider cells within are CD4* and high for CD25* heave a suppressive cepacity in vitro (25). We therefore purified CD4* Tr cells from peripheral blood, and by FACS sorting, purified CD25****

(25). We therefore purified CD4* Tr cells from peripheral blood, and by FACS sorting, purified CD25****

(25) and CD25*****

(26) provided the control of the con

[0035] As one of the defining characteristics of CD25*CD4+ Tr cells is constitutive and high expression of CD25, we screened the CD25*CD4+ T-cell clones for expression of CD25 at least 10 days after restimulation (in the resting phase). A shown in Figure 6, the clones displayed a heterogeneous expression of CD25. Approximately half the clones remained 99-100% positive for CD25, and possessed a relatively high mean fluorescence intensity (MFI). Other clones contained a significant number of CD25* cells and a lower MFI. T-cell clones derived the CD25*CD4+ T cells displayed a low excendance of CD25* cells in the resting phase and consequently also a low MFI.

[0036] The clones were subsequently fested for their shillip to proliferate in response to activation via the TOR in the absence or presence of exogenous IL-2. It has been well established that both murine and human CD25*CD4*T cells fall to proliferate in response to acD3 mAbs in the absence of costimulation via CD28 and/or addition of IL-2 (23, 24, 26). In order to determine if this was also true at the clonal level, we tested the CD25*CD4*T-cell clones for their ability to proliferate in the presence or absence of IL-2. Similar to the heterogeneity observed in terms of prospession of CD25, the clones were also heterogeneous in terms of proliferated well in the presence of IL-2. The remaining clones (1472, 20%) proliferated well in response to acD3 mAbs even in the absence of IL-2. The remaining clones (1472, 20%) proliferated well in response to acD3 mAbs even in the absence of IL-2. A representative subset of the 72 cloned tested is shown in Figure 7. In contrast, amongst the CD25*CD4*T-cell clones tested, the majority (14/22, 65%) proliferated well in response to acD3 mAbs alone.

[0037] The heterogeneity of CD25*CD4* T-cell clones in terms of expression of CD25 and proliferation suggested that even within the 0.6% CD25fn⁴CD4* Tr cell population of PBMCs, not all cells may be suppressor cells, and that a proportion could be activated T helper cells. To test this hypothesis we performed in vitro suppression assays. As shown in Figure 8, indeed only a subset of the CD25*CD4* T cell clones were able to suppress the proliferative response of autologous CD4* T cells in response to a CD3* of mbs cross-infect on T-cell-depleted PBMCs (8A) or immobilized on

plastic (8B). Interestingly, only those clones which were anergic and displayed a constitutively high expression of CD25 had a suppressive phenotype. When data from all the CD25+CD4+ T-cell clones tested were pooled together, and the clones were separated into suppressive and non-suppressive groups, expression of CD25 was found to be significantly higher in the group with suppressive activity (p<0.000007) (Table 2). In contrast, proliferation in response to aCD3 mAbs was a less reliable predictor of suppressive capacity, as several clones within the non-suppressive category were anergic. These data indicate that constitutively high expression of CD25 is a marker for CD4+ T regulatory cells at the clonal level. [0038] We also determined the cytokine production profile of the CD25+CD4+T-cell clones. As shown in Table 3, nonsuppressive clones tended to possess a Th0 pattern of cytokine production and produced moderate levels of most cytokines tested. In contrast, all CD25+CD4+T-cell clones which had suppressive activity produced significant amounts of TGF-B, small and variable amounts of IL-4, IL-5 and IFN-y, but failed to produce detectable levels of IL-2 or IL-10, These data indicate that CD25*CD4* T regulatory cells are likely distinct from IL-10-producing Tr1 cells, although it cannot be excluded that they represent the same cells at different stages of differentiation. The fact that the only cytokine which was consistently detected in the supernatants of all the suppressive CD25+CD4+T-cell clones was TGF-β suggests that they are more likely related to the TGF-8 producing Th3 cells which were originally described in models of oral tolerance (27, 28).

TABLES

[0039]

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Table 1. Cytokine production by CD25+CD4+ Tr cells. Purified cells were stimulated as indicated and supernatants were collected after 24 (for IL-2) or 72 hours. The amount of cytokine was determined by ELISA. Data represent the average values (pc/ml) of pooled data from 4 independent experiments. Cytokine production by allogeneic APCs alone has been subtracted.

25	Stimuli	Cells	IL-2	IL-4	IL-10	IFN-γ	TGF-β	
	αCD3/28 +IL-2	CD25+	N.D.	153	1148	5723	1322	
	αCD3/28 +IL-2	CD25	N.D.	94	840	9773	1225	
	allogeneic APCs	CD25+	<20	<20	298	527	509	
30	allogeneic APCs	CD25	99.5	<20	251	5744	637	

Table 2. Suppressive CD25+CD4+ T-cell clones have a distinct phenotype from non-suppressive clones. Summary of the phenotype of all the CD25+CD4+ T-cell clones which were extensively characterized. Percent suppression represents the average reduction of proliferation of autologous CD4+ T cells upon activation with αCD3 mAbs. immobilized on plastic or T-cell-depleted ACPs, in the presence of the indicated T-cell clones, Numbers represent the average suppression observed in 2-6 independent experiments. MFI represents the average expression of CD25 as determined in 2-6 independent tests, cpm represents the amount of thymidine incorporated following activation with αCD3 mAbs immobilized on plastic. Numbers represent the average of duplicates in a single test, and are representative of results obtained in several subsequent tests.

	representative or results out	nive of results obtained in several subsequent tests.			
		suppression (%)	CD25 MFI	αCD3 (cpm)	
	non-suppressive				
	2	0	36	15947	
45	3	0	83	166	
	6	0	40	908	
	37	0	40	36280	
	84	n.t.	19	20281	
50	85	0	33	1839	
	86	0	91	23139	
	87	0	21	4166	
	88	n.t.	34	36376	
	89	0	43	17702	
55	90	0	46	21210	
	92	0	33	34004	
	03	n t	10	20005	

EP 1 409 650 B1

Table continued

		suppression (%)	CD25 MFI	αCD3 (cpm)
	94	n.t.	12	6929
5	95	0	66	410
	suppressive			
	4	37	90	244
10	12	40	100	122
	13	52	101	511
	15	24	211	644
	17	36	110	88
	18	73	97	123
15	19	60	88	1289
	20	22	98	285
	21	76	65	72
	22	39	102	180
	24	45	174	281
	28	45	179	380
20	29	54	113	98
	40	63	85	108
	42	45	77	52
25	47	57	86	61
	48	56	68	66
	57	31	130	600

Nt: not tested

Table 3. Cytokine production profile of CD25⁺CD4⁺ T cell clones. T-cell clones were activated with αCD3 and αCD28 mAbs, and supernatants were collected after 24 (for IL-2) or 48 hours. Amounts of cytokines in the supernatants were determined by capture ELISA and/ or CBA assay as described in the materials and methods. n.t.; not tested.

			IL-4	IL-5	IL-10	IFN-y	TGF-β
15		(pg/ml)	(pg/ml)	(ng/ml)	(pg/ml)	(ng/ml)	(pg/ml)
	non-suppressive						
	2	<20	1184	35.4	79	2.7	181
	3	<20	8002	3.5	98	0.8	257
20	6	<20	57	0.9	<20	0.1	94
	.37	56	552	3.9	17	7.7	n.t.
	84	540	607	4.5	39	7.0	7024
	85	<20	521	7.9	94	1.5	214
25	86	394	537	3.9	62	12.0	n.t.
	87	<20	419	3.0	125	4.2	70
	88	<20	168	2.5	<20	1.1	n.t.
	89	476	604	4.3	101	8.8	n.t.
30	90	1360	618	4.6	100	12.6	n.t.
	92	280	1963	10.2	163	6.7	217
	95	<20	46	0.4	169	0.5	329
35	Suppressive						l I
	4	<20	<20	<0.02	<20	<0.06	31
	17	<20	67	0.05	<20	0.07	277
	18	<20	<20	<0.02	<20	n.t.	141
40	19	<20	140	0.3	<20	0.1	311
	20	<20	<20	<0.02	<20	<0.06	44
	21	<20	<20	<20	<20	<0.06	130
	22	<20	<10	<0.02	<20	<0.06	410
45	29	<20	<20	<0.02	<20	<0.06	182
	40	<20	<20	<0.02	<20	<0.06	206
	42	<20	83	0.2	<20	0.2	369
	57	<20	<10	<0.02	<20	0.1	278

MATERIALS AND METHODS

^{1.} Isolation and characterization of human CD25+CD4+ Tr cells

^{50 [040]} Purification of CD25*CD4*T reells, Human periphenal blood was obtained from healthy donors in accordance with local eithical committee approval. PBMCs were prepared by centrifugation over Fool-Hypauge gradents (Nycomed Amerikam, Uppseia, Sweden), and CD4*T cells were purified by positive or negative selection (by depletion of CD8, CD1*tb, CD16, CD19, CD38 and CD58 boositive cells with the CD4* Mullisork fix of the Unitoxide CD4*T cell Isolation.

kit, respectively (Miltenyi Biotech, Gladbach, Germany), Following isolation of CD4+T cells, CD25* cells were stained with PE-coupled anti-CD25 mAbs and purified following addition of anti-PE coupled magnetic beads (Miltenyi Biotech), Alternatively, CD4+T cells were purified with magnetic beads directly-coupled to anti-CD25 (Miltenyi Biotech) to facilitate FACS analysis. Results ottained with CD25+CD4+T cells solated by negative or positive selection, and directly or indirectly coupled CD25 mAbs were identical. Staining with 21/0* PBMCs, typically 2-2x1/0* CD25+CD4+T cells were isolated, with a purity ranging from 90-95%. CD25+CD4+T cells were also collected, with a purity ranging from 70-95%. For purification of CD25+cells following in vitra activation of CD25+cells (CD25+CD4+T cells were calvated of 48 hours with immobilized anti-CD3 (10pg/mi) and soluble anti-CD28 (1pg/mi) mAbs and CD25+T cells were purified as described

(9041] In vitro expansion of T cell fines. CD25*CD4* To CD25*CD4* To cells were isolated as described. To cells (2x105 cellsmit) were stimulated with anti-CD3 († µg/mi) (OKT3, Orthoclone, Jansen Clag, Italy) in the presence of an allogeneic feeder-cell mixture consisting of 1x105*PGMCs/ml, (irradiated 6000 RADS) and 1x105*JY cellsmit (irradiated 10,000 RADS), an EBV-LCL which expresses high levels of HLA and costimulatory molecules as well as cytokines, as previously described (23,90.). All cultures were performed in X-Vivo 15 medium (BloWhitakev, Begramo, Italy) supplemented with 10% FCS (Mascia Brunelli, Milan, Italy), 1% pooled human serum, 100 Um1 penicillin/streptomycin (Bristol-Myers Squibb, Sermoneta, Italy) and 2mM glutamine (GibcoBRL, Milan, Italy) (are referred to as complete medium). Three days after activation, 40Uml ril c.2 (Chrion Italia), Milan, Italy) was added. Cells were split as necessary and fresh medium with IL-2 was added. T-cell lines were restimulated every 14 days. All experiments on expanded cells were performed at least 10 days after activation.

20 [0042] Proliferation and suppression of Toells. To analyze proliferation in response to polyclonal activation, 98 well round-bottom piates (Costar) were costed overnight at 4°C with anti-CD3 mAbs (10,µg/ml) in 0.1M Tris, pH 9.5, and washed three times with PBS. T cells were plated at an initial density of 5x10° cells/ml (100,000 cells/well) in a final volume of 200µl of complete medium in the absence or presence of soluble anti-CD28 mAbs (1µg/ml) (Pharmingen, San Diego, CA), soluble secondary rabbit anti-nuces Abs (10,µg/ml) (Signa, Millar, Ilayl) and/or IL-2 (100U/ml).

[0043] To test antigen-specific Toell profiferation, freshly isolated CD25*CD4* Tr or CD25*CD4* T cells (2.5x10* cells/ml) were stimulated with irradiated (6003 Rebs) allogeneic PBMCs (2.5x10* cells/ml) that had been depleted of CD3* cells by negative selection (Dynal, Oxold). For suppression, increasing numbers (up to 2.5x10* cells/ml) of freshly isolated autologous CD25*CD4* Tr cells were added. Cells were co-cultured in a final volume of 200µl of complete medium in 98 well round-bottom plates. Control 196 (10ap/ml) (1967), Pharmigen), naturizing anti-1-10f (10ap/ml) (1967), Pharmigen) and/or anti-1CF-5, p.23 (10ap/ml or 50ap/ml) (180.D), or F(ab²), anti-CTLA+ (10ap/ml) (Ancell, Bayport, MN) mAbs were added as indicated. For suppression of memory T cells, cells from expanded CD25*CD4* T-cell lines were cultured with allogeneic APCs (from a donor different from that used in the allogeneic feeder-cell mixture), and increasing numbers of expanded autologous CD25*CD4* T cells were added as described above. For control experiments, CD25*CD4**T cells uprefiled from In vitro-activated CD25*CD4** Toells were added in increasing numbers to freshly numbers to freshly antibody of the control experiments.

[0044] After the indicated time, wells were pulsed for 16 hours with 1µCi/well 3H-thymidine (Amersham, Uppsala, Sweden). Cells were harvested, and counted in a scintillation counter.

[0045] ELISAs. For detection of IL-10, IL-4, IL-2, IFN-y, and TGF-β, capture ELISAs were performed on supernatants of cells (1x10⁵ Tcells/ml) that had been stimulated with immobilized anti-CD3 mAbs (10₄g/ml) with or without anti-CD28 (1₄g/ml) and IL-2 (100U/ml), or irradiated CD3-elepted allogenes (PMICs (1x10° cells/ml)) or 24 (0x10° cells/ml) or 24 (0x10° c

[0046] FACS analysis. Anti-CD4, -CD25, -HLA-DR, -CD45RD, -CD821, -CD68 and -CD401. mAbs were purchased from Backton Dekinson (Mountain New, CA) and were directly coupled to FTIC or PE. Expression of IL-2PB (CD122) and IL-2

2. Isolation and characterization of human CD25+CD4+ Tr cell clones

isolated autologous CD25 CD4+ T cells and allogeneic APCs.

[0047] Purification and cloning of CD25+CD4+ Tr cells. CD4+T cells from PBMCs were obtained as described above.

CD4** Tealls were stained with FTIC-coupled anti-CD4 and PE-coupled anti-CD25 mAbs (Becklon-Dickson) and CD25* and CD25* cells were purified by FACS-sorting on a FACStar (Becklon-Dickson), CD25*CD4* and CD25* CD4** Teals were subsequently cloned at 1 cell/well in 36*-well round bottom plates by limiting dilution in the presence of an allogeneic feader-cell mixture consisting of 5x105* PBMCs/ml, 5x105* JY cells/ml and 0.05µg/ml PHA. After 3 days, IL-2 (40 U/ml) was added. Teal clones were cultivare in X-vivo 18 with 55*. Human Serum. At day 14, goving wells were picked and re-stimulated with an allogeneic feeder-cell mixture as described above. Clones were split as necessary, and restimulated as above every 14 days. The medium was replenished every 3-5 days. Clones were used for experiments between days 10 and 14 of the previous re-stimulation (i.e. in the resting phase).

[0048] Proliferation and suppression of T cells. To analyze the proliferative capacity of T-cell clones in response to polycional activation, 98 will usuad-bottom plates (Costar) were coated overnight at 4"C with anti-CD3 mAbs (10g, 4m) in 0.1M Tris, p.H 9.5, and washed three times with PBS. T-cell clones were plated at an initial density of 2x10⁵ cells/ml (40,000 cells/well) as finally oldume of 200µl of complete medium in the absence or presence of IL-2 (100U/ml). To test for the capacity of T-cell clones to suppress the proliferation of autologous CD4*T cells, fresh CD4*T cells were purified by positive selection (Miltery) Biotech) and stimulated with anti-CD3 mAbs which had been immobilized on plastic (as 5 described above) or bound to allogenable CD3*Ceplated PBMSc (freadlated 6000 RADS). CD4*T-cells (40,000 cells/well)

described above) or bound to allogeneic CD3-depleted PBMCs (irradiated 6000 RADS). CD4+T cells (40,000 cells/well) were cultured alone, or in the presence of a 1:1 ratio of T-cell clones in a final volume of 200µl of complete medium in 96 well round-bottom plates.

[0049] After the indicated time, wells were pulsed for 16 hours with 1µCi/well 3H-thymidine (Amersham, Uppsala, Sweden). Cells were harvested, and counted in a scintillation counter.

20 [050] EL/SAs, T Coll clones (1x10⁶ collexin) were stimulated with immobilized anti-CD3 mAbs (10,μg/ml) and anti-CD28 (1μg/ml), and supernatants were collected after 24 hours for IL-2 and after 48 hours for IL-2 in the collected of GF-β in additional source and the collected after 24 hours for IL-2 in Late 1 (1 to 1 to 1 and IR-N-ywere after determined of their by capture ELISA as described above. Lavels of IL-2, IL-4, IL-5, IL-1 to and IR-N-ywere after determined either by capture ELISA (ID Blosciences) as described above or by the cytometric bead array kit (CBA) (ID Blosciences), according to the manufacture's instructions. A direct comparison of capture 25 ELISA and CBA demonstrated that the two methods were highly comparable in terms of the amount of cytokine detected in the suscendant.

[0051] Statistical Analysis. All analysis for statistically significant differences were performed with the student's paired ttest, p values less than 0.05 were considered significant. Results are expressed as means ± SEM.

30 Abbreviations

[0052]

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CD cluster of differentiation

IL interleukin

TGF transforming growth factor

APC antigen presenting cell

mAb monoclonal antibody
FACS fluorescence activated cell sorting

PHA phytohemoagalutinin

PBMC peripheral blood mononuclear cells

MFI mean fluorescence intensity

EBV Ebstein-Barr virus

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Claims

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- The use of ex-vivo isolated and expanded human CD25+CD4+ Tr cells for the preparation of immunomodulating or immunosuppressive agents.
- The use of ex-vivo isolated human CD25+CD4+ Tricell clones constitutively expressing CD25 for the preparation of immunomodulating or immunosuppressive agents.
 - The use according to claim 1 or 2, for the prevention or therapy of graft-vs-host disease, organ rejection, autoimmune diseases and for the prevention of adverse immune responses to transgenes and vector-derived proteins after gene therapy.
 - The use according to claim 1, wherein human CD25*CD4* Tr cells are expanded in vitro under one or more of the following conditions: co-culture with feeder-cell mixture, polycional stimulation, antigen specific stimulation, addition of cytokine.
- The use according to claim 2, wherein the CD25+CD4+ Tr cell clones constitutively expressing CD25 are isolated ex-vivo by the following steps:
 - a) purifying CD4+T cells from PBMCs;
 - b) separating CD25+ from CD25-T cells:
 - c) cloning CD25+CD4+ T cells by limiting dilution;
 - d) stimulation with phytohemagglutinin or and-CD3 mAb, in the presence of IL-2;
 - e) selecting the suppressive clones that display a constitutively high expression of CD25.
 - An immunosuppressive agent containing ex-vivo expanded human CD25+CD4+Tr cells or isolated CD25+CD4+Tr cell clones constitutively expressing CD25.
 - 7. An immunosuppressive agent according to claim 6, further containing cytokines or additional immunosuppressants.
- An immunosuppressive agent according to claims 6-7, which is in form of stabilized cell preparation.
 - 9. A method of isolating immunosuppressive CD25+CD4+ Tr cell clones which comprises the steps of:
 - a) purifying CD4*T cells from PBMCs;
- b) separating CD25+ from CD25- T cells;
 - c) cloning CD25+CD4+ T cells by limiting dilution;
 - d) stimulation with phytohemagglutinin or anti-CD3 mAb, in the presence of IL-2;
 - c) selecting the suppressive clones that display a constitutively high expression of CD25.
 - 10. A method according to claim 9, wherein the stimulation according to step (d) is carried out in the presence of an allogenic or autologous feeder-cell mixture consisting of irradiated PBMCs.
 - 11. A method according to claim 10, which is carried out with irradiated autologous or allogeneic EBV-transformed cell lines.
- 12. A method according to claim 9, wherein in step (d) the suppressive clones are selected on the basis of the following characteristics:
 - 100% constant-positivity for CD25 expression in the resting phase at least 10 days after stimulation with

- phytohemagglutinin or anti-CD3 mAb in the presence of IL-2;
- expression of CD25 at a significantly higher level in comparison to T cell clones isolated in parallel from CD25 CD4+ T cells or non suppressive clones isolated from CD25+CD4+ T cells.
- 5 13. Isolated CD25+CD4+ Tr cell clones obtainable by the process of claims 9-12.
 - Isolated CD25+CD4+ Tricell clones according to claim 13, which do not produce IL-2.
- 15. The use of CD25*CD4* Tr cell clones according to claims 13-14 for the preparation of in vitro systems for the identification of molecules that modulate the immune response.
 - 16. The use according to claim 15, in large scale gene expression arrays, differential proteomies screenings and for the generation of monoclonal antibodies

15 Patentansprüche

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- Verwendung von ex vivo isolierten und expandierten humanen CD25*CD4*-Tr-Zeilen zur Herstellung von Immunmodulatoren oder Immunsuporessiva.
- Verwendung von ex vivo isolierten humanen CD25+CD4+-Tr-Zellklonen, welche konstitutiv CD25 exprimieren, zur Herstellung von Immunmodulatoren oder Immunsuppressiva.
- Verwendung nach Anspruch 1 oder 2 zur Verh
 ütung der Therapie von Transplantat-Wirt-Abstoßung, Organabsto ßung, Autoimmunerkrankungen und zur Verh
 ütung negativer immunreaktionen auf Transgene und Vektor-abgeleitete Proteine nach einer Gentherapie.
- Verwendung nach Anspruch 1, wobei humane CD25¹CD4¹-Tr-Zellen in vitrounter einer oder mehreren der folgenden Bedingungen expandiert werden: Cokultur mit einer Feederzell-Mischung, polyklonale Stimulation antigenspezifisehe Stimulation, Zugabe von Zytokinen.
 - Verwendung nach Anspruch 2, wobei die CD25+CD4+-Tr-Zellklone, die konstitutiv CD25 exprimieren, ex vivo mit den folgenden Schritten isoliert werden:
 - a) Reinigung von CD4+-T-Zellen aus PBMCs;
 - b) Abtrennen von CD25+- von CD25-T-Zellen:
 - c) Klonierung von CD25+CD4+-T-Zellen durch Grenzverdünnung;
 - d) Stimulation mit Phytohämagglutinin oder Anti-CD3-mAb, in Gegenwart von IL-2;
 - e) Selektion der suppressiven Klone, welche eine konstitutiv hohe Expression von CD25 zeigen.
 - Immunsuppressivum, enthaltend ex vivo expandierte humane CD25+CD4+-Tr-Zellen oder isolierte CD25+CD4+-Tr-Zellklone, die konstitutiv CD25 exprimieren.
 - 7. Immunsuppressivum nach Anspruch 6, ferner enthaltend Zytokine oder zusätzliche Immunsuppressivu.
 - 8. Immunsuppressivum nach den Ansprüchen 6-7, welches in Form einer stabilisierten Zellpräparation vorliegt.
 - 9. Verfahren zur Isolierung von immunsuppressiven CD25*CD4*-Tr-Zellklonen, umfassend die Schritte:
 - a) Reinigung von CD4+-T-Zellen aus PBMCs;
 - b) Abtrennen von CD25+- von CD25-T-Zellen:
 - c) Klonierung von CD25+CD4+-T-Zellen durch Grenzverdünnung;
 - d) Stimulation mit Phytohämagglutinin oder Anti-CD3-mAb, in Gegenwart von IL-2;
 - e) Selektion der suppressiven Klone, welche eine konstitutiv hohe Expression von CD25 zeigen.
 - Verfahren nach Anspruch 9, wobei die Stimulation gemäß Schritt (d) in Gegenwart einer allogenen oder autologen Feederzell-Mischung, bestehend aus bestrahlten PBMCs, durchgeführt wird.

- Verfahren nach Anspruch 10, welches mit bestrahlten autologen oder allogenen EBV-transformierten Zeillinien durchgeführt wird.
- 12. Verfahren nach Anspruch 9, wobei in Schritt (d) die suppressiven Klone auf Basis der folgenden charakteristischen Merkmale ausgewählt werden:
 - 100 % konstante Positivität für CD25-Expression in der Ruhephase mindestens 10 Tage nach Stimulation mit Phytohämagglutinin oder Anti-CD3-mAb in Gegenwart von IL-2;
- Expression von CD25 auf einem signifikant h\u00f6heren Niveau im Vergleich zu T-Zellklonen, die parallel aus CD25-CD4+T-Zeilen isoliert wurden, oder nicht-suppressiven Klonen, die aus CD25+CD4+T-Zeilen isoliert wurden.
 - Isolierte CD25+CD4+-Tr-Zellklone, erhältlich mit dem Verfahren der Ansprüche 9 12
- 14. Isolierte CD25+CD4+-Tr- Zellkione nach Anspruch 13, welche kein IL-2 produzieren.
 - Verwendung von CD25+CD4+-Tr-Zellklonen nach den Ansprüchen 13-14 zur Herstellung von in vitro-Systemen für die Identifizierung von Molekülen, welche die Immunreaktion modulieren.
- 16. Verwendung nach Anspruch 15 in Genexpressions-Arrays in großem Maßstab, differentiellen proteonomischen Screenings und zur Herstellung monoklonaler Antikörper.

Revendications

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- Utilisation de cellules Tr CD25+CD4+ humaines isolées ex vivo et multipliées pour la préparation d'agents immunomodulateurs ou immunosuporessifs.
- Utilisation de clones de cellules Tr CD25*CD4* humaines isolés ex vivo, exprimant de manière constitutive CD25
 pour la préparation d'agents immunomodulateurs ou immunosuppressifs.
 - 3. Utilisation selon la revendication 1 ou 2, pour la prévention ou la thérapie de la maladie du greffon contre l'hôte, du rejet d'organe, des maladies auto-immunes ou pour la prévention des réponses immunitaires néfastes contre les transpènes et les protéliers détriées de vectuurs après une thérapie génique.
 - 4. Utilisation seion la revendication 1, dans laquelle les cellules Tr CD25+CD4+ humaines sont multipliées in vitro dans une ou plusieurs des conditions suivantes: co-culture avec un mélange de cellules nourricières, stimulation poéycionale, stimulation spédifique d'artipène, addition de cytokines.
- 5. Utilisation selon la revendication 2, dans laquelle les clones de cellules Tr CD25*CD4* exprimant de manière constitutive CD25 sont isolés ex vivo suivant les étapes suivantes:
 - a) purification des cellules T CD4+ à partir de PBMC;
 - b) séparation de CD25+ des cellules T CD25-;
 - c) clonage des cellules T CD25+CD4+ par dilution limitante;
 - d) stimulation avec de la phytohémagglutinine ou du mAb anti-CD3, en présence de IL-2;
 - e) sélection des clones suppressifs qui présentent une expression constitutivement élevée de CD25.
 - Agent immunosuppressif contenant des cellules Tr CD25+CD4+ humaines multipliées ex vivo ou des clones de cellules Tr CD25+CD4+ isolés, exprimant de manière constitutive CD25.
 - Agent immunosuppressif selon la revendication 6, contenant en outre des cytokines ou des immunosuppresseurs supplémentaires.
- Agent immunosuppressif selon les revendications 6 et 7, qui est sous forme de préparation cellulaire stabilisée.
 - 9. Procédé d'isolement de clones de cellules Tr CD25+CD4+ immunosuppressifs, qui comprend les étapes de:

- a) purification des cellules T CD4+ à partir de PBMC;
- b) séparation de CD25+ des cellules T CD25-;

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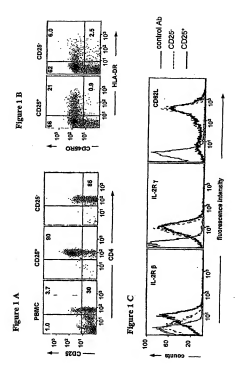
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- c) clonage des cellules T CD25+CD4+ par dilution limitante;
- d) stimulation avec de la phytohémagglutinine ou du mAb anti-CD3, en présence de IL-2:
- e) sélection des clones suppressifs qui présentent une expression constitutivement élevée de CD25.
 - 10. Procédé selon la revendication 9, dans lequel la stimulation selon l'étape (d) est réalisée en présence d'un mélange de cellules nourricières allogéniques ou autologues constitué de PBMC irradiées.
- 10 11. Procédé selon la revendication 10, qui est mis en oeuvre avec des lignées de cellules transformées par EBV, autologues ou allogéniques irradiées,
 - 12. Procédé selon la revendication 9, dans lequel dans l'étape (d), les clones suppressifs sont choisis sur la base des caractéristiques suivantes:
 - un taux de positivité constant de 100% de l'expression de CD25 dans la phase de repos au moins 10 jours après la stimulation avec la phytohémagolutinine ou le mAb anti-CD3 en présence de IL-2;
 - une expression de CD25 à un niveau nettement supérieur par rapport à des ciones de cellules T isolés en parallèle à partir de cellules T CD25CD4+ ou des clones non suppressits isolés à partir de cellules T CD25CD4+.
 - 13. Clones de cellules Tr CD25+CD4+ isolés, pouvant être obtenus par le procédé des revendications 9 à 12.
 - 14. Clones de cellules Tr CD25+CD4+ isolés selon la revendication 13, qui ne produisent pas de IL-2.
- 25 15. Utilisation de clones de cellules Tr CD25+CD4+ selon les revendications 13 et 14 pour la préparation de systèmes in vitro pour l'identification de molécules qui modulent la réponse immunitaire.
 - 16. Utilisation selon la revendication 15, dans des panoplies d'expression génique à grande échelle, des criblages par protéomique différentielle et pour la production d'anticorps monoclonaux.



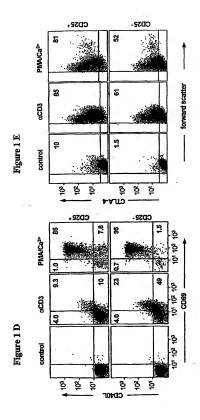


Figure 2

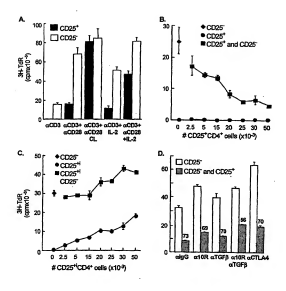
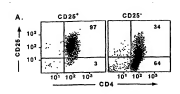
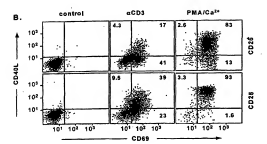


Figure 3





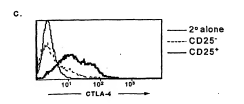


Figure 4

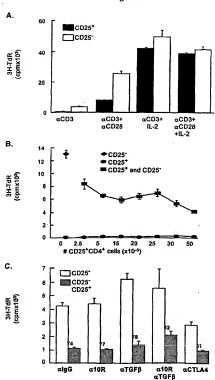
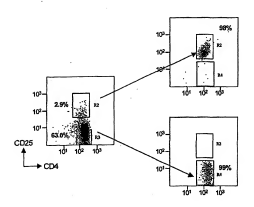
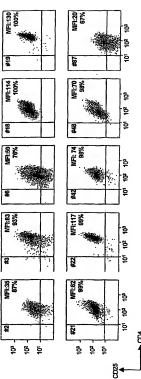


Figure 5







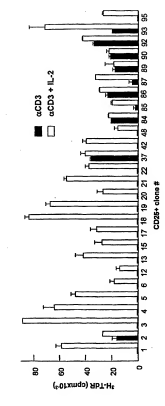


Figure 7

Figure 8

